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LOGINID:SSPTACDR1614

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TERMINAL (ENTER 1, 2, 3, OR ?):2

* * * * * * * * * Welcome to STN International * * * * * * * * *

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NEWS 14 MAR 31 IFICDB, IFIPAT, and IFIUDB enhanced with new custom IPC display formats
NEWS 15 MAR 31 CAS REGISTRY enhanced with additional experimental spectra
NEWS 16 MAR 31 CA/CAplus and CASREACT patent number format for U.S. applications updated
NEWS 17 MAR 31 LPCI now available as a replacement to LDPCI
NEWS 18 MAR 31 EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS 19 APR 04 STN AnaVist, Version 1, to be discontinued
NEWS 20 APR 15 WPIDS, WPINDEX, and WPIX enhanced with new predefined hit display formats
NEWS 21 APR 28 EMBASE Controlled Term thesaurus enhanced
NEWS 22 APR 28 IMSRESEARCH reloaded with enhancements
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NEWS 25 JUN 06 EPFULL enhanced with 260,000 English abstracts
NEWS 26 JUN 06 KOREAPAT updated with 41,000 documents
NEWS 27 JUN 13 USPATFULL and USPAT2 updated with 11-character patent numbers for U.S. applications
NEWS 28 JUN 19 CAS REGISTRY includes selected substances from web-based collections
NEWS 29 JUN 25 CA/CAplus and USPAT databases updated with IPC reclassification data
NEWS 30 JUN 30 AEROSPACE enhanced with more than 1 million U.S. patent records
NEWS 31 JUN 30 EMBASE, EMBAL, and LEMBASE updated with additional options to display authors and affiliated

NEWS 32 JUN 30 STN on the Web enhanced with new STN AnaVist
Assistant and BLAST plug-in
NEWS 33 JUN 30 STN AnaVist enhanced with database content from EPFULL

NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3,
AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.

NEWS HOURS STN Operating Hours Plus Help Desk Availability
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Enter NEWS followed by the item number or name to see news on that specific topic.

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FILE 'HOME' ENTERED AT 11:46:39 ON 16 JUL 2008

```
=>
Uploading
THIS COMMAND NOT AVAILABLE IN THE CURRENT FILE
Do you want to switch to the Registry File?
Choice (Y/n):
Switching to the Registry File...
Some commands only work in certain files. For
command can only be used to look at the index
index. Enter "HELP COMMANDS" at an arrow prom
commands which can be used in this file.
```

=> FILE REGISTRY

COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST	0.21	0.21

FILE 'REGISTRY' ENTERED AT 11:46:59 ON 16 JUL 2008
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
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STRUCTURE FILE UPDATES: 15 JUL 2008 HIGHEST RN 1034171-01-1

New CAS Information Use Policies, enter HELP USAGETERMS for details

TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

Please note that search-term pricing does apply when using the search function.

conducting SmartSELECT searches.

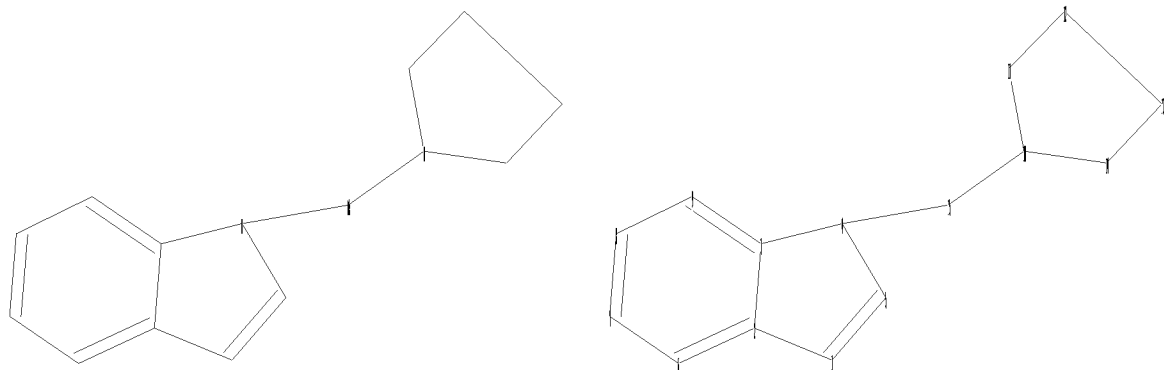
predicted properties as well as tags indicating availability of

experimental property data in the original document. For information on property searching in REGISTRY, refer to:

<http://www.cas.org/support/stngen/stndoc/properties.html>

=>

Uploading C:\Program Files\STNEXP\Queries\10566094a1.str



chain nodes :

15

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14

chain bonds :

4-15 10-15

ring bonds :

1-2 1-5 2-3 2-6 3-4 3-9 4-5 6-7 7-8 8-9 10-14 10-11 11-12 12-13 13-14

exact/norm bonds :

1-2 1-5 3-4 4-5 4-15 10-14 10-11 10-15 11-12 12-13 13-14

normalized bonds :

2-3 2-6 3-9 6-7 7-8 8-9

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom

11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS

Element Count :

Node 15: Limited

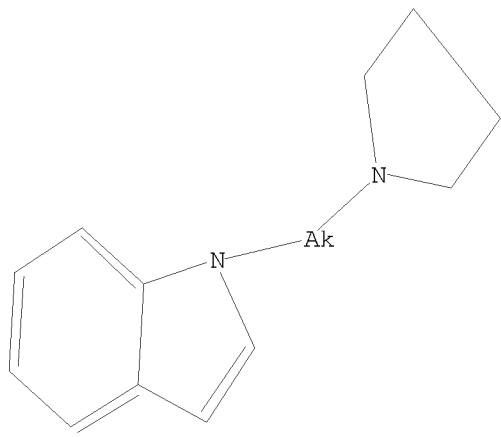
C,C1-3

L1 STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS

L1 STR



Structure attributes must be viewed using STN Express query preparation.

```
=> s l1 sss sam
SAMPLE SEARCH INITIATED 11:47:12 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 103791 TO ITERATE
```

```
1.9% PROCESSED      2000 ITERATIONS          3 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
```

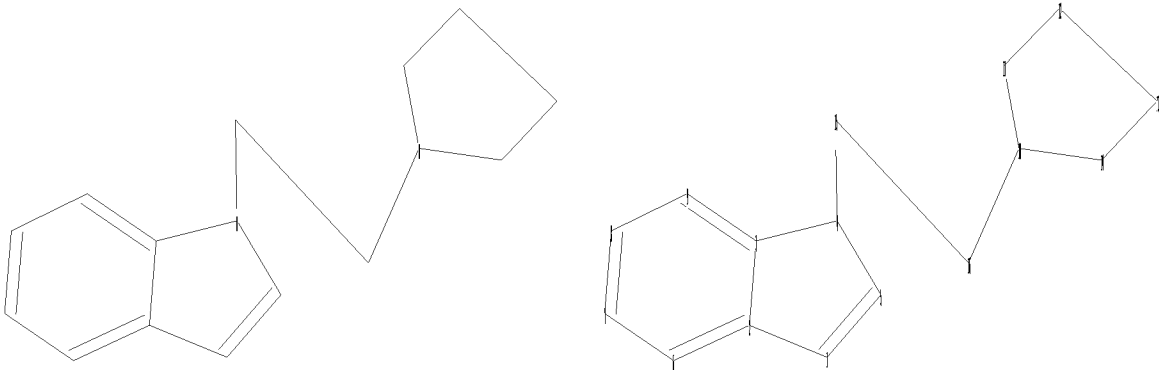
```
FULL FILE PROJECTIONS: ONLINE  **INCOMPLETE**
                        BATCH   **INCOMPLETE**
PROJECTED ITERATIONS:    2056698 TO 2094942
PROJECTED ANSWERS:        2365 TO     3861
```

L2 3 SEA SSS SAM L1

```
=> s l1 sss fam
'SSS' IS NOT VALID HERE
For additional help, enter "HELP SEARCH".
```

```
=> s l1 fam sam
STRUCTURES CONTAINING VARIABLE NODES NOT VALID IN EXACT OR FAMILY SEARCH
You have requested a full structure (EXA or FAM) search on a
structure containing one of the special variable-atom symbols
A, M, Q, or X, or a variable group G. Only bond variability
is allowed in structures for EXA or FAM searches. Variable
nodes are never permitted.
```

```
=>
Uploading C:\Program Files\STNEXP\Queries\10566094a2.str
```



chain nodes :

15 16

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14

chain bonds :

4-15 10-16 15-16

ring bonds :

1-2 1-5 2-3 2-6 3-4 3-9 4-5 6-7 7-8 8-9 10-14 10-11 11-12 12-13 13-14

exact/norm bonds :

1-2 1-5 3-4 4-5 4-15 10-14 10-11 10-16 11-12 12-13 13-14

exact bonds :

15-16

normalized bonds :

2-3 2-6 3-9 6-7 7-8 8-9

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom
11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS 16:CLASS

L3 STRUCTURE UPLOADED

=> s 13 sss sam

SAMPLE SEARCH INITIATED 11:48:44 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 8480 TO ITERATE

23.6% PROCESSED 2000 ITERATIONS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

16 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 164080 TO 175120

PROJECTED ANSWERS: 862 TO 1850

L4 16 SEA SSS SAM L3

=> s 13 sss full

FULL SEARCH INITIATED 11:48:58 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 167937 TO ITERATE

100.0% PROCESSED 167937 ITERATIONS

1330 ANSWERS

SEARCH TIME: 00.00.01

L5 1330 SEA SSS FUL L3

=> file caplus
COST IN U.S. DOLLARS SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 179.74 179.95

FILE 'CAPLUS' ENTERED AT 11:49:06 ON 16 JUL 2008
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FILE COVERS 1907 - 16 Jul 2008 VOL 149 ISS 3
FILE LAST UPDATED: 15 Jul 2008 (20080715/ED)

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=> s 15
L6 211 L5

=> file reg
COST IN U.S. DOLLARS SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 0.48 180.43

FILE 'REGISTRY' ENTERED AT 11:49:40 ON 16 JUL 2008
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DICTIONARY FILE UPDATES: 15 JUL 2008 HIGHEST RN 1034171-01-1

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<http://www.cas.org/support/stngen/stndoc/properties.html>

=> s 16 and serotonin
528 SEROTONIN
L7 0 L6 AND SEROTONIN

=> s 16 and 5ht
18 5HT
L8 0 L6 AND 5HT

=> s 16
L9 1330 L5

=> s 16 and 5-ht
15225065 5
2519 HT
66 HTS
2580 HT
(HT OR HTS)
295 5-HT
(5(W)HT)
L10 0 L6 AND 5-HT

=> s 16 and appetite
6 APPETITE
L11 0 L6 AND APPETITE

	SINCE FILE ENTRY	TOTAL SESSION
COST IN U.S. DOLLARS		
FULL ESTIMATED COST	28.05	208.48

FILE 'CAPLUS' ENTERED AT 11:52:39 ON 16 JUL 2008
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=>

```
=> d ibib abs hitstr 200
L11 HAS NO ANSWERS
'IBIB ABS HITSTR ' IS NOT A VALID STRUCTURE FORMAT KEYWORD
Structure Formats
SIA ----- Structure Image, Attributes, and map table if it contains
        data. (Default)
SIM ----- Structure IMage.
SAT ----- Structure ATtributes and map table if it contains data.
SCT ----- Structure Connection Table and map table if it contains
        data.
SDA ----- All Structure DAta (image, attributes, connection table and
        map table if it contains data).
NOS ----- NO Structure data.
ENTER STRUCTURE FORMAT (SIA), SCT, SDA, SIM, SAT, NOS:end
```

=> s 15

L12 211 L5

=> d ibib abs hitstr 200

```
L12 ANSWER 200 OF 211 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 1966:19187 CAPLUS
DOCUMENT NUMBER: 64:19187
ORIGINAL REFERENCE NO.: 64:3492g-h,3493a
TITLE: 11-(Tertiary aminoalkyl)-5,6-dihydro-11H-
benzo[a]carbazoles
INVENTOR(S): Freed, Meier E.; Rice, Leonard M.; Hertz, Elisabeth
PATENT ASSIGNEE(S): American Home Products Corp.
SOURCE: 3 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Unavailable
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
```

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
-----	-----	-----	-----	-----
US 3215691		19651102	US 1962-168505	19620124
PRIORITY APPLN. INFO.:			US	19620124

AB The title compds. are claimed to have antidepressant, anorectic, and anti-histaminic activity. 5,6-Dihydro-11H-benzo[a]carbazole (11 g.) and 1.44 g. NaH in 100 ml. HCONMe₂ was stirred 1 hr. at 35-40°, treated with 6 g. Me₂N(CH₂)₃Cl and the mixture stirred overnight. The mixture was cooled, poured into ice-H₂O, acidified with dilute HCl, extracted with Et₂O, and the aqueous layer basified with NaOH. The oily product which separated was extracted with Et₂O and the Et₂O extract was washed with aqueous NaCl, dried,

and

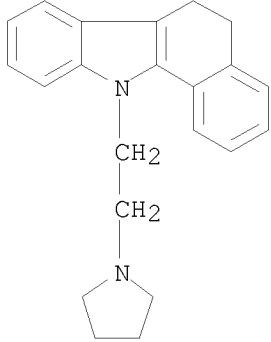
evaporated to give 11-(3-dimethylaminopropyl)-5,6-dihydro-11H-benzo[a]carbazole, m. 59-60° (90% EtOH) [HCl salt m. 222-3° (Me₂CO-MeOH)]. Similarly prepared were the following substituted 5,6-dihydro-11H-benzo[a]carbazoles (substituent given): 11-(2-piperidinoethyl), m. 91-2° (HCl salt m. 214-222°; fumarate salt m. 204-6°); 11-(2-dimethylaminoethyl), (HCl salt m. 228-31°); 11-(2-pyrrolidinoethyl), m. 85.6-86.0° (fumarate salt m. 192.5-3°); 11-(2-dimethylaminoethyl), b0.08 168° (fumarate salt m. 147-50°); 11-(2-dimethylaminopropyl), (fumarate salt m. 159-62°); 11-(2-morpholinoethyl), m. 122-3°; 11-(3-dipropylaminopropyl) [fumarate salt m. 142-4° (decomposition)]; and 11-(6-dimethylaminohexyl) (fumarate salt m. 104-7°).

IT 4624-82-2P, 5H-Benzo[a]carbazole, 6,11-dihydro-11-[2-(1-pyrrolidinyl)ethyl]- 4624-83-3P, 5H-Benzo[a]carbazole, 6,11-dihydro-11-[2-(1-pyrrolidinyl)ethyl]-, fumarate (1:1)

RL: PREP (Preparation)
(preparation of)

RN 4624-82-2 CAPLUS

CN 5H-Benzo[a]carbazole, 6,11-dihydro-11-[2-(1-pyrrolidinyl)ethyl]- (CA INDEX NAME)



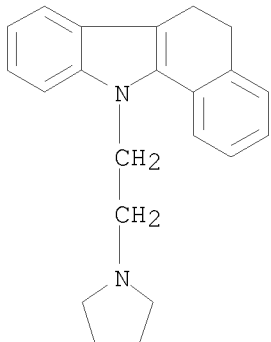
RN 4624-83-3 CAPLUS

CN 5H-Benzo[a]carbazole, 6,11-dihydro-11-[2-(1-pyrrolidinyl)ethyl]-, fumarate (1:1) (8CI) (CA INDEX NAME)

CM 1

CRN 4624-82-2

CMF C22 H24 N2



CM 2

CRN 110-17-8

CMF C4 H4 O4

Double bond geometry as shown.



```
=> end
ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF
LOGOFF? (Y)/N/HOLD:logoff
'LOGOFF' IS NOT VALID HERE
For an explanation, enter "HELP LOGOFF".
```

```
=> file reg
COST IN U.S. DOLLARS          SINCE FILE      TOTAL
                                ENTRY           SESSION
FULL ESTIMATED COST          8.33            216.81
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE      TOTAL
                                                ENTRY           SESSION
CA SUBSCRIBER PRICE          -0.80            -0.80
```

```
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DICTIONARY FILE UPDATES: 15 JUL 2008 HIGHEST RN 1034171-01-1

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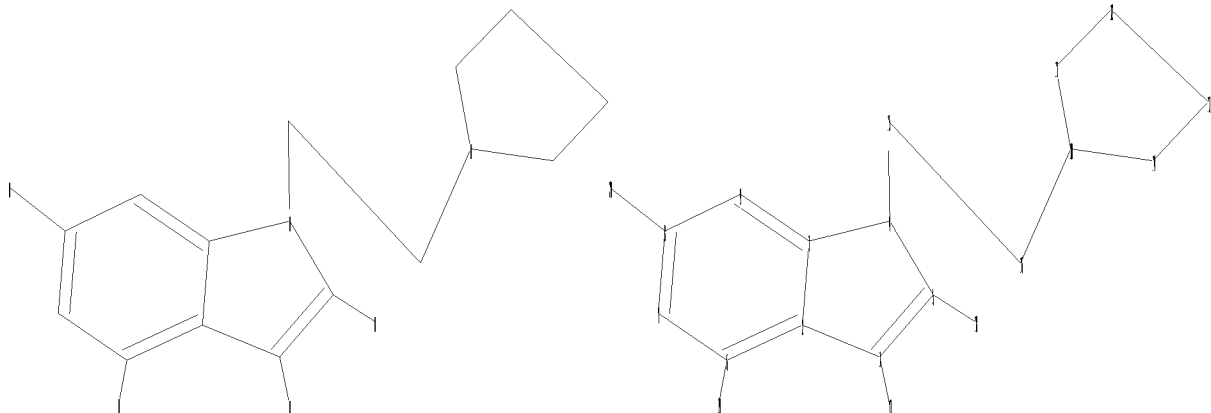
TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

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<http://www.cas.org/support/stngen/stndoc/properties.html>

```
=>
Uploading C:\Program Files\STNEXP\Queries\10566094a3.str
```



```
chain nodes :
15 16 17 18 19 20
ring nodes :
1 2 3 4 5 6 7 8 9 10 11 12 13 14
```

```

chain bonds :
1-17 4-15 5-18 6-19 8-20 10-16 15-16
ring bonds :
1-2 1-5 2-3 2-6 3-4 3-9 4-5 6-7 7-8 8-9 10-14 10-11 11-12 12-13 13-14

exact/norm bonds :
1-2 1-5 3-4 4-5 4-15 10-14 10-11 10-16 11-12 12-13 13-14
exact bonds :
1-17 5-18 6-19 8-20 15-16
normalized bonds :
2-3 2-6 3-9 6-7 7-8 8-9

```

```

Match level :
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom
11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS 16:CLASS 17:CLASS 18:CLASS
19:CLASS 20:CLASS

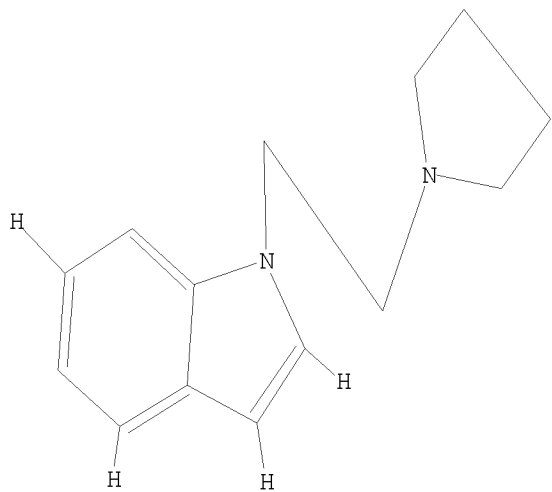
```

L13 STRUCTURE UPLOADED

```

=> d 113
L13 HAS NO ANSWERS
L13 STR

```



Structure attributes must be viewed using STN Express query preparation.

```

=> s 113 sss sam
SAMPLE SEARCH INITIATED 11:56:32 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 8480 TO ITERATE

```

```

23.6% PROCESSED 2000 ITERATIONS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
3 ANSWERS

```

```

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**
PROJECTED ITERATIONS: 164080 TO 175120
PROJECTED ANSWERS: 41 TO 467

```

L14 3 SEA SSS SAM L13

=> s 113 sss full
FULL SEARCH INITIATED 11:56:37 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 167937 TO ITERATE

100.0% PROCESSED 167937 ITERATIONS 86 ANSWERS
SEARCH TIME: 00.00.01

L15 86 SEA SSS FUL L13

=> file caplus
COST IN U.S. DOLLARS SINCE FILE TOTAL
 ENTRY SESSION
FULL ESTIMATED COST 178.36 395.17

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL
 ENTRY SESSION
CA SUBSCRIBER PRICE 0.00 -0.80

FILE 'CAPLUS' ENTERED AT 11:56:42 ON 16 JUL 2008
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FILE COVERS 1907 - 16 Jul 2008 VOL 149 ISS 3
FILE LAST UPDATED: 15 Jul 2008 (20080715/ED)

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=> s 115
L16 27 L15

=> d ibib abs hitstr 15

L16 ANSWER 15 OF 27 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2004:725572 CAPLUS
DOCUMENT NUMBER: 142:211383
TITLE: Medicinal Chemistry Driven Approaches Toward Novel and Selective Serotonin 5-HT6 Receptor Ligands
AUTHOR(S): Holenz, Joerg; Merce, Ramon; Diaz, Jose Luis; Guitart, Xavier; Codony, Xavier; Dordal, Alberto; Romero, Gonzalo; Torrens, Antoni; Mas, Josep; Andaluz, Blas; Hernandez, Susana; Monroy, Xavier; Sanchez, Elisabeth; Hernandez, Enrique; Perez, Raquel; Cubi, Roger;

CORPORATE SOURCE: Sanfeliu, Olga; Buschmann, Helmut
 Departments of Medicinal Chemistry, Discovery Biology
 and Discovery Chemistry, Laboratorios Dr. Esteve S.A.,
 Barcelona, 08041, Spain
 SOURCE: Journal of Medicinal Chemistry (2005), 48(6),
 1781-1795
 CODEN: JMCMAR; ISSN: 0022-2623
 PUBLISHER: American Chemical Society
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 142:211383

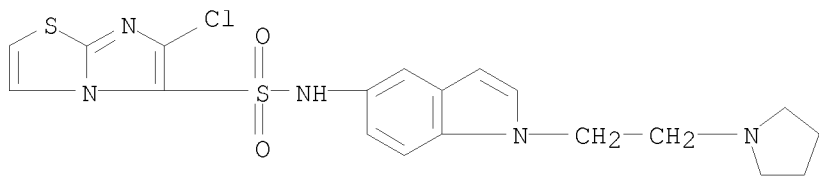
AB Based on a medicinal chemical guided hypothetical pharmacophore model, novel series of indolyl sulfonamides have been designed and prepared as selective and high-affinity serotonin 5-HT₆ receptor ligands. Furthermore, based on a screening approach of a discovery library, a series of benzoxazinepiperidinyl sulfonamides were identified as selective 5-HT₆ ligands. Many of the compds. described in this paper possess excellent affinities, displaying pKi values greater than 8 (some even >9) and high selectivities against a wide range (>50) of other CNS relevant receptors. First, structure-affinity relationships of these ligands are discussed. In terms of functionality, high-affinity antagonists, as well as agonists and even partial agonists, were prepared. Compds. 19c and 19g represent the highest-affinity 5-HT₆ agonists ever reported in the literature. These valuable tool compds. should allow for the detailed study of the role of the 5-HT₆ receptor in relevant animal models of disorders such as cognition deficits, depression, anxiety, or obesity.

IT 753020-85-8P

RL: DMA (Drug mechanism of action); PAC (Pharmacological activity); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
 (medicinal chemical driven approaches toward novel and selective serotonin 5-HT₆ receptor ligands)

RN 753020-85-8 CAPLUS

CN Imidazo[2,1-b]thiazole-5-sulfonamide, 6-chloro-N-[1-[2-(1-pyrrolidinyl)ethyl]-1H-indol-5-yl]- (CA INDEX NAME)

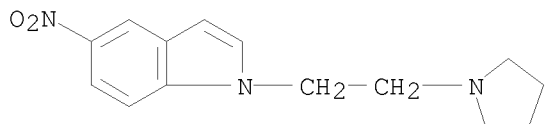


IT 753021-17-9P 753021-23-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (medicinal chemical driven approaches toward novel and selective serotonin 5-HT₆ receptor ligands)

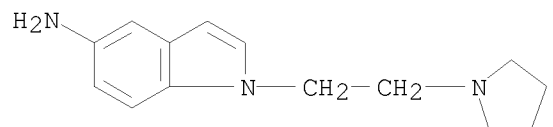
RN 753021-17-9 CAPLUS

CN 1H-Indole, 5-nitro-1-[2-(1-pyrrolidinyl)ethyl]- (CA INDEX NAME)



RN 753021-23-7 CAPLUS

CN 1H-Indol-5-amine, 1-[2-(1-pyrrolidinyl)ethyl]- (CA INDEX NAME)



REFERENCE COUNT: 68 THERE ARE 68 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=>